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## WHAT IS CLAIMED IS:

- 1. A method for inducing a protective mucosal cytotoxic T lymphocyte (CTL) response in a mammalian subject comprising contacting a mucosal tissue of the subject with a composition comprising a purified soluble antigen.
- The method of claim 1, wherein the soluble antigen is an antigenic peptide.
- 3. The method of claim 1, wherein said composition further comprises an adjuvant.
- The method of claim 3, wherein the adjuvant is selected from cholera toxin (CT), mutant cholera toxin (MCT),
   or mutant- E. coli heat labile enterotoxin (MLT).
  - 5. The method of claim 1, further comprising administering a purified cytokine to the subject.
- 6. The method of claim 1, wherein the cytokine is contacted with a mucosal surface of the subject.
- 7. The method of claim 5, wherein the purified cytokine is selected from granulocyte-macrophage colonystimulating factor (GM-CSF), interleukin-2 (IL-2), interleukin-7 (IL-7), interleukin-12 (IL-12) or tumor necrosis factor a (TNFa).
- The method of claim 1, further comprising
   administering purified interferon-γ to the subject.
  - 9. The method of claim 8, wherein the purified interferon- $\gamma$  is contacted with a mucosal surface of the subject.
  - 10. The method of claim 5, further comprising administering purified interferon- $\gamma$  to the subject.

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- 11. The method of claim 10, wherein the purified interferon- $\gamma$  is contacted with a mucosal surface of the subject.
- 12. The method of claim 1, wherein said composition further comprises a purified cytokine selected from granulocyte-macrophage colony-stimulating factor (GM-CSF), interleukin-2 (IL-2), interleukin-7 (IL-7), interleukin-12 (IL-12) or tumor necrosis factor.
- 13. The method of claim 1, wherein said composition further comprises purified interferon- $\gamma$ .
- 14. The method of claim 12, wherein said composition15 further comprises purified interferon-γ.
  - 15. The method of claim 1, wherein the antigen is a peptide derived from a pathogenic virus.
- 20 16. The method of claim 15, wherein the pathogenic virus is HIV-1.
  - 17. The method of claim 15, wherein the pathogenic virus is influenza virus.
  - 18. The method of claim 15, wherein the pathogenic virus is rotavirus.
- 19. The method of claim 1, wherein the antigen is a peptide derived from a pathogenic bacterium or protozoan.
  - 20. The method of claim 1, wherein the antigen is a tumor-associated peptide.
- 21. The method of claim 1, wherein the antigen is a peptide comprising an HIV-1 cluster peptide vaccine construct (CLUVAC) selected from the group consisting of: EQMHEDIISLWDQSLKPCVKRIQRGPGRAFVTIGK (SEQ ID NO:1),

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KQIINMWQEVGKAMYAPPISGQIRRIQRGPGRAFVTIGK (SEQ ID NO:2),
RDNWRSELYKYKVVKIEPLGVAPTRIQRGPGRAFVTIGK (SEQ ID NO:3),
AVAEGTDRVIEVVQGAYRAIRHIPRRIRQGLERRIQRGPGRAFVTIGK (SEQ ID NO:4), DRVIEVVQGAYRAIRHIPRRIRQGLERRIQRGPGRAFVTIGK (SEQ ID NO:5), DRVIEVVQGAYRAIRRIQRGPGRAFVTIGK (SEQ ID NO:6),
AQGAYRAIRHIPRRIRRIQRGPGRAFVTIGK (SEQ ID NO:7),
EQMHEDIISLWDQSLKPCVKRIHIGPGRAFYTTKN (SEQ ID NO:8),
KQIINMWQEVGKAMYAPPISGQIRRIHIGPGRAFYTTKN (SEQ ID NO:9),
RDNWRSELYKYKVVKIEPLGVAPTRIHIGPGRAFYTTKN (SEQ ID NO:10),
AVAEGTDRVIEVVQGAYRAIRHIPRRIRQGLERRIHIGPGRAFYTTKN (SEQ ID NO:11), DRVIEVVQGAYRAIRHIPRRIRQGLERRIHIGPGRAFYTTKN (SEQ ID NO:12), DRVIEVVQGAYRAIRRIHIGPGRAFYTTKN (SEQ ID NO:13) and
AQGAYRAIRHIPRRIRRIHIGPGRAFYTTKN (SEQ ID NO:14).

- 15 22. The method of claim 21, wherein the HIV-1 CLUVAC is HIV-1 CLUVAC PCLUS3-18IIIB (SEQ ID NO:2).
  - $23\,.$  The method of claim 21, wherein the HIV-1 CLUVAC is HIV-1 CLUVAC PCLUS3-18MN (SEQ ID NO:9).
  - 24. The method of claim 21, wherein the HIV-1 CLUVAC is HIV-1 CLUVAC PCLUS 6.1-18MN (SEQ ID NO:12).
- 25. A method for inducing a protective mucosal CTL response in a subject, comprising contacting a mucosal tissue of the subject with a composition comprising a soluble antigen, wherein said composition does not comprise an adjuvant.
- 26. The method of claim 25, further comprising administering a purified cytokine to the subject.
  - 27. The method of claim 25, wherein the cytokine is contacted with a mucosal surface of the subject.
  - 28. The method of claim 27, wherein the purified cytokine is selected from granulocyte-macrophage colony-stimulating factor (GM-CSF), interleukin-2 (IL-2),

interleukin-7 (IL-7), interleukin-12 (IL-12) or tumor necrosis factor a (TNFa).

- 29. The method of claim 25, further comprising administering purified interferon-γ to the subject.
  - 30. The method of claim 29, wherein the purified interferon- $\gamma$  is contacted with a mucosal surface of the subject.

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- 31. The method of claim 26, further comprising administering purified interferon- $\gamma$  to the subject.
- 32. The method of claim 31, wherein the purified interferon-γ is contacted with a mucosal surface of the subject.
- 33. The method of claim 25, wherein said composition further comprises a purified cytokine selected from granulocyte-macrophage colony-stimulating factor (GM-CSF), interleukin-2 (IL-2), interleukin-7 (IL-7), interleukin-12 (IL-12) or tumor necrosis factor.
- 34. The method of claim 25, wherein said composition25 further comprises purified interferon-γ.
  - 35. The method of claim 33, wherein said composition further comprises purified interferon- $\gamma$ .
  - 36. The method of claim 25, wherein the antigen is a peptide derived from a pathogenic virus.
  - 37. The method of claim 36, wherein the pathogenic virus is HIV-1.

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38. The method of claim 36, wherein the pathogenic virus is influenza virus.

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- 39. The method of claim 36, wherein the pathogenic virus is rotavirus.
- 40. The method of claim 25, wherein the antigen is a peptide derived from a pathogenic bacterium or protozoan.
  - 41. The method of claim 25, wherein the antigen is a tumor-associated peptide.
- 10 The method of claim 25, wherein the antigen is a speptide comprising an HIV-1 cluster peptide vaccine construct (CLUVAC) selected from the group consisting of: EQMHEDIISLWDQSLKPCVKRIQRGPGRAFVTIGK (SEQ ID NO:1), KQIINMWQEVGKAMYAPPISGQIRRIORGPGRAFVTIGK (SEQ ID NO:2), 15 RDNWRSELYKYKVVKIEPLGVAPTRIORGPGRAFVTIGK (SEQ ID NO:3), AVAEGTDRVIEVVQGAYRAIRHIPRRIRQGLERRIQRGPGRAFVTIGK (SEQ ID NO:4), DRVIEVVQGAYRAIRHIPRRIRQGLERRIQRGPGRAFVTIGK (SEQ ID NO:5), DRVIEVVQGAYRAIRRIORGPGRAFVTIGK (SEQ ID NO:6), AQGAYRAIRHIPRRIRRIQRGPGRAFVTIGK (SEQ ID NO:7), 20 EQMHEDIISLWDQSLKPCVKRIHIGPGRAFYTTKN (SEQ ID NO:8), KQIINMWQEVGKAMYAPPISGQIRRIHIGPGRAFYTTKN (SEQ ID NO:9), RDNWRSELYKYKVVKIEPLGVAPTRIHIGPGRAFYTTKN (SEQ ID NO:10), AVAEGTDRVIEVVQGAYRAIRHIPRRIRQGLERRIHIGPGRAFYTTKN (SEQ ID
  - 43. The method of claim 42, wherein the HIV-1 CLUVAC is HIV-1 CLUVAC PCLUS3-18IIIB (SEQ ID NO:2).

NO:11), DRVIEVVQGAYRAIRHIPRRIRQGLERRIHIGPGRAFYTTKN (SEQ ID

NO:12) \_\_DRVIEVVQGAYRAIRRIHIGPGRAFYTTKN (SEQ ID NO:13) and

AQGAYRAIRHIPRRIRRIHIGPGRAFYTTKN (SEQ ID NO:14).

- 44. The method of claim 42, wherein the HIV-1 CLUVAC is HIV-1 CLUVAC PCLUS3-18MN (SEQ ID NO:9).
- 45. The method of claim 42, wherein the HIV-1 CLUVAC is HIV-1 CLUVAC PCLUS 6.1-18MN (SEQ ID NO:12).
  - 46. An immunogenic composition for inducing a protective mucosal CTL response in a subject and adapted for

intrarectal administration comprising a purified soluble antigen formulated for intrarectal delivery to the rectum, colon, sigmoid colon, or distal colon.

- 5 47. The immunogenic composition of claim 46, which comprises a rectal enema, foam, suppository, or topical gel.
- 48. The immunogenic composition of claim 46, further comprising a base, carrier, or aabsorption-promoting agent adapted for intrarectal delivery.
  - 49. The immunogenic composition of claim 48, which includes a rectal emulsion or gel preparation.
- 15 50. The immunogenic composition of claim 48, wherein the soluble antigen is admixed with a homogenous gel carrier.
  - 51. The immunogenic composition of claim 48, wherein the homogenous gel carrier is a polyoxyethylene gel.
  - 52. The immunogenic composition of claim 48, wherein the soluble antigen is admixed with a rectally-compatible foam.
- 25 \_\_53. The immunogenic composition of claim 48, wherein the soluble antigen is formulated in a suppository.
  - 54. The immunogenic composition of claim 53, wherein the suppository is comprised of a base selected from a polyethyleneglycol, witepsol H15, witepsol W35, witepsol E85, propyleneglycol dicaprylate (Sefsol 228), Miglyol810, hydroxypropylcellulose-H (HPC), or carbopol-934P (CP).
- 55. The immunogenic composition of claim 53, comprising at least two base materials.

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- 56. The immunogenic composition of claim 46, including a stabilizing agent to minimize intrarectal degradation of the soluble antigen.
- 5 57. The immunogenic composition of claim 46, including an absorption-promoting agent.
  - 58. The immunogenic composition of claim 57, wherein the absorption-promoting agent is selected from a surfactant, mixed micelle, enamines, nitric oxide donor, sodium salicylate, glycerol ester of acetoacetic acid, clyclodextrin or beta-cyclodextrin derivative, or medium-chain fatty acid.
- 59. The immunogenic composition of claim 46, further comprising an adjuvant.
  - 60. The immunogenic composition of claim 59, wherein the adjuvant is selected from cholera toxin (CT), mutant cholera toxin (MCT), mutant- E. coli heat labile enterotoxin, or pertussis toxin.
  - 61. The immunogenic composition of claim 59, wherein the adjuvant is conjugated to a mucosal tissue or T cell binding agent.
  - 62. The immunogenic composition of claim 61, wherein the mucosal tissue or T cell binding agent is selected from protein A, an antibody that binds a mucosal tissue- or T-cell-specific protein, or a ligand or peptide that binds a mucosal tissue- or T-cell-specific protein.
  - 63. The immunogenic composition of claim 59, wherein the adjuvant comprises a recombinant cholera toxin (CT) having a B chain of CT substituted by protein A conjugated to a CT A chain to eliminate toxicity and enhance mucosal tissue binding mediated by protein A.

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- 64. The immunogenic composition of claim 59, wherein the adjuvant is conjugated to a protein or peptide that binds specifically to T cells.
- 5 65. The immunogenic composition of claim 64, wherein the protein or peptide binds to CD4 or CD8.
  - 66. The immunogenic composition of claim 66, wherein the protein or peptide is an HIV V3 loop or T cell-binding peptide fragment thereof.
  - 67. The immunogenic composition of claim 59, further comprising purified IL-12.
- 15 68. The immunogenic composition of claim 59, further comprising purified interferon-γ.
  - 69. The immunogenic composition of claim 68, further comprising purified IL-12.